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Molecular signatures of plastic phenotypes in two eusocial insect species with simple societies

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Phenotypic plasticity is important in adaptation and shapes the evolution of organisms. However, we understand little about what aspects of the genome are important in facilitating plasticity. Eusocial insect societies produce plastic phenotypes from the same genome, as reproductives (queens) and nonreproductives (workers). The greatest plasticity is found in the simple eusocial insect societies in which individuals retain the ability to switch between reproductive and nonreproductive phenotypes as adults. We lack comprehensive data on the molecular basis of plastic phenotypes. Here, we sequenced genomes, microRNAs (miRNAs), and multiple transcriptomes and methylomes from individual brains in a wasp (*Polistes canadensis*) and an ant (*Dinoponera quadricaps*) that live in simple eusocial societies. In both species, we found few differences between phenotypes at the transcriptional level, with little functional specialization, and no evidence that phenotype-specific gene expression is driven by DNA methylation or miRNAs. Instead, phenotypic differentiation was defined more subtly by nonrandom transcriptional network organization, with roles in these networks for both conserved and taxon-restricted genes. The general lack of highly methylated regions or methylome patterning in both species may be an important mechanism for achieving plasticity among phenotypes during adulthood. These findings define previously unidentified hypotheses on the genomic processes that facilitate plasticity and suggest that the molecular hallmarks of social behavior are likely to differ with the level of social complexity.

social evolution | phenotypic plasticity | genome sequencing | transcriptomes | DNA methylation

Phenotypic plasticity allows organisms to maintain fitness in a changing environment. Plasticity influences organismal ecological resilience, adaptability, evolutionary innovations, and speciation (1, 2). However, we understand little about the molecular signatures (the genes involved and differential regulation thereof) of such plasticity. Determining the molecular basis of phenotypic plasticity is fundamental to our understanding of the building blocks of life and has the potential to uncover insights into selection for adaptive function and phenotypic innovation (3–5).

The profound action of evolution in the generation of biological diversity can be discerned from the genome (6). However, genome sequence alone is not sufficient to explain diverse phenotypic variation because such analyses infer associations based on gene evolution and gene sharing rather than directly identifying

differentially expressed genes (DEGs) in the phenotypes of interest (7). Here, in addition to genome and microRNA (miRNA) sequencing, we use deep transcriptome and methylome sequencing of single brains from alternative phenotypes to determine the differential molecular processes associated with highly plastic phenotypes in two species of eusocial insects (8).

Significance

In eusocial insect societies, such as ants and some bees and wasps, phenotypes are highly plastic, generating alternative phenotypes (queens and workers) from the same genome. The greatest plasticity is found in simple insect societies, in which individuals can switch between phenotypes as adults. The genomic, transcriptional, and epigenetic underpinnings of such plasticity are largely unknown. In contrast to the complex societies of the honeybee, we find that simple insect societies lack distinct transcriptional differentiation between phenotypes and coherently patterned DNA methylomes. Instead, alternative phenotypes are largely defined by subtle transcriptional network organization. These traits may facilitate genomic plasticity. These insights and resources will stimulate new approaches and hypotheses that will help to unravel the genomic processes that create phenotypic plasticity.

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Conflict of interest statement: S.B. is a founder and shareholder of Cambridge Epigenetix Limited, and W.R. is a consultant and shareholder of Cambridge Epigenetix Limited.

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Data deposition: Genomic analyses were performed on the whole-genome assemblies of *Polistes canadensis* and *Dinoponera quadricaps*, deposited at the DNA Data Bank of Japan/European Molecular Biology Laboratory/GenBank under the accession nos. PRJNA253269 and PRJNA253275, respectively. Raw data from all bisulfite-sequencing and RNA-sequencing libraries were deposited in the Gene Expression Omnibus (GEO) database (accession no. GSE59525).

See Commentary on page 13755.

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Hymenopteran eusocial insects exhibit enormous interspecific variation in phenotypic plasticity, in the form of reproductive (queen) and nonreproductive (worker) phenotypes (9), across multiple independent origins (10). Our two study species (the dinosaur ant *Dinoponera quadricaps* and the paper wasp *Polistes canadensis*) exhibit very simple societies, where individuals retain the ability to switch phenotype (11, 12). This characteristic contrasts with the adult honey bee *Apis mellifera* and most ants, which exhibit low levels of phenotypic plasticity and have been the focus of most previous molecular analyses (13). Our two study species share similar levels of plasticity among individuals, with a single reproductive egg-layer (“gamergate” in *D. quadricaps* and “queen” in *P. canadensis*) that is morphologically identical to the nonreproductives; if the reproductive dies, it is quickly replaced by one of the nonreproductives. Both species share many ecological traits but evolved social phenotypes independently (14, 15) (Dataset S1). As such, we present two independent studies on the molecular basis of highly plastic phenotypes in these simple societies (Fig. 1A and B).

Our aims were threefold. First, we sequenced the genomes of *P. canadensis* and *D. quadricaps* to provide genomic baseline data for eusocial insect species with simple societies, including the first aculeate wasp genome sequence. Second, we sequenced and analyzed individual brain transcriptomes to identify differential transcription patterns associated with phenotypes. Third, we sequenced global miRNAs and individual-level phenotype-specific brain methylomes to determine the extent to which these putative regulators associate with phenotypic differentiation and genomic organization. These analyses highlight fundamental traits of the molecular basis of phenotypic differentiation and plasticity of similar phenotypes apparent in both species. As such, these data provide the first genome sequence, to our knowledge, for an aculeate wasp; provide a framework and hypotheses for revealing the molecular signatures of caste evolution; and, more generally, help define scenarios where conserved or contrasting molecular processes in phenotypic evolution might be used.

Results and Discussion

Typical Insect Genome Composition and Organization. A single haploid male for each species was sequenced on the Illumina platform achieving 110-fold coverage. The de novo assembled *P. canadensis* and *D. quadricaps* genomes were 211 Mega-basepairs (Mbp) and 268 Mbp in size, respectively (SI Text, sections I, II.1, and II.2). These genome sequences are almost complete, with 97–99% of the conserved cluster of orthologous proteins mapped in the two genomes; 79–86% of proteins were annotated (Fig. S1A–D and SI Text, sections II.3–II.5). The genome compositions were similar to

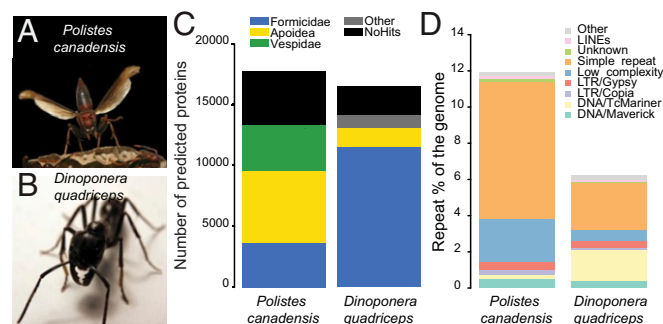


Fig. 1. Genome sequencing and organization. *P. canadensis* (A) and *D. quadricaps* (B) share similar ecological, social, and behavioral traits (Dataset S1). (C) *P. canadensis* shares more similarity in predicted proteins with bees (Apidae) than ants (Formicidae), as expected, given the lack of other published aculeate wasp genome sequences; *D. quadricaps* shares greatest similarity of predicted protein sequences with sequenced ant genomes (Formicidae). These data are derived from computational protein analyses (SI Text, section II.7). (D) Distribution of different classes of repetitive elements and transposons across *P. canadensis* and *D. quadricaps* genomes.

the genome sequences of other social insects, with *D. quadricaps* sharing more of its predicted protein content with other ants (Formicidae), whereas *P. canadensis* shows more equitable levels of protein sharing with ants (Formicidae) and bees (Apidae) (Fig. 1C; Fig. S1E; SI Text, section II.6; and Dataset S1). This difference is likely to reflect the absence of any other aculeate wasp genome sequence in the public domain. Finally, the genome of *P. canadensis* contains more transposable elements (452,247, 12% of the genome) than *D. quadricaps* (217,417, 6% of the genome), most of which are simple or low-complexity repeats (Fig. 1D and SI Text, section II.7). Transposable elements were recently identified as potentially important in the evolution of social complexity in bees (6).

Low Levels of Transcriptional Differentiation Between Phenotypes.

We obtained over 100 gigabase pairs (Gbps) of brain transcriptome sequence data from 23 individual adult female brains (four to seven biological replicates each of reproductives and nonreproductives per species), generating, on average, 3.6 Mbp (20.29 ± 0.67 -fold coverage) and 4.9 Mbp (17.4 ± 1.36 -fold coverage) per individual for the wasp and ant, respectively (SI Text, sections III.1 and III.2 and Dataset S2). In both species, we found fewer than 1% of genes were differentially expressed (DEGs), with little evidence of functional specialization between phenotypes (5). Using the union of DEGs from EdgeR [parametric approach (16)] and NOISeq [nonparametric approach (17)] (Fig. 2; Table 1; SI Text, section III.3; and Dataset S2), we found 67 (0.4%) DEGs in *P. canadensis* and 147 (0.8%) DEGs in *D. quadricaps*. In both species, the nonparametric approach identified significantly more up-regulated genes in reproductives relative to nonreproductives ($\chi^2 = 31$, $P = 2.2e-08$; Table 1). In *P. canadensis*, gene expression in nonreproductives was found to be more stochastic (noisy) than in reproductives despite similar variance of expression among the biological replicates (Fig. S2). Recent research suggests that evolution can shape noise in gene expression and that such noise can be adaptive and heritable (18–20). If noise in transcription is an indicator of phenotypic plasticity (21–23), our results would suggest that transcription in the non-reproductive phenotype is more responsive to changes in the biotic and social environment than transcription in the reproductive phenotype. Despite the small number of DEGs, significant functional enrichment of DEGs was detected in the ant reproductives, with 29 gene ontology terms significantly enriched for functions that included metabolic and ribosomal processes, regulation of expression, and an extracellular component [false discovery rate (FDR) < 0.5; SI Text, section III.4 and Dataset S2]. There was little sign of functional enrichment in the wasp (5) (although before FDR correction, oxidoreductase activity and lipid transport were overrepresented in reproductives). These data suggest there is little phenotypic specialization in the brain tissue of either species.

No Distinct Methylation Patterning Across the Genome or Between Phenotypes.

We sequenced the methylomes from three biological replicates each of individual adult brains from reproductive and nonreproductive phenotypes in *P. canadensis* and *D. quadricaps* using whole-genome bisulfite sequencing [BS-seq; 20 gigabase (GB) (>10-fold coverage) per brain] (SI Text, section IV.1 and Dataset S3).

We compared methylation patterns with the honey bee (24) to provide a reference point because the honey bee is the only close relative to our study species with comparable data on brain methylation available (SI Text, section IV.2). Global levels of methylation in the cytosine-guanine (CG) context were similar in both species, and similar to the honey bee (Table 2). *P. canadensis* exhibited greater methylation in the non-CG context but significantly fewer highly methylated regions than *D. quadricaps* (Table 2; Fig. S3A and B; and SI Text, section IV.3). However, in comparison to the honey bee, both species showed relatively little gene body-specific methylation targeting (Fig. 3A; Table 2; Fig. S3C; and SI Text, section IV.4), together with a striking lack of consistently fully methylated cytosines (Fig. 3B). In both *P. canadensis* and *D. quadricaps*, DNA methylation is dispersed sparsely across genes (Fig. 3C), particularly

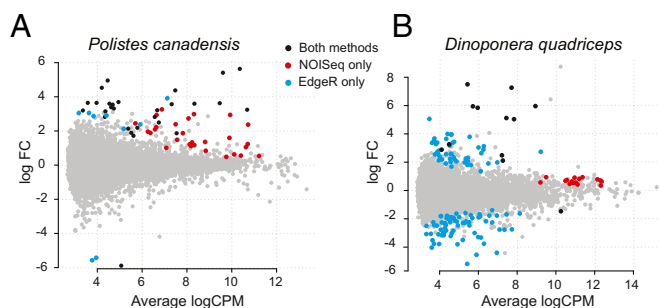


Fig. 2. Low levels of transcriptional differentiation between phenotypes. (A and B) Counts per million plots of log fold mean gene expression differences between phenotypes, showing the numbers and log fold differences of DEGs up-regulated in reproductives (positive) and nonreproductives (negative). The union and individual results of two methods for detecting DEGs (NOISeq and EdgeR) are presented. FC, fold change.

in *P. canadensis*, whose genome lacks a DNA methyltransferase 3 (DNMT3) gene, an enzyme involved in de novo methylation (25, 26) (Fig. S4A and S5A and SI Text, section IV.5). In *P. canadensis*, we also found a prevalence of asymmetrical (one strand only) CG methylation, together with a variant of the DNA methylation transferase 1 (DNMT1) gene involved in the maintenance of DNA methylation, in its genome (Fig. S4 B–D and SI Text, section IV.5). As observed in *A. mellifera* brains (27), both study species possess and express a ten-eleven translocation (TET) hydroxylase gene and base excision repair genes involved in demethylation, and have detectable hydroxymethylation in brain tissue (Figs. S4F and S5 and SI Text, section IV.6). Together, these general features provide an epigenetic landscape that may facilitate plasticity of genome function.

Despite the general paucity of methylation patterning, we found significant conservation of methylated orthologs (Fig. 3D; SI Text, section IV.7; and Dataset S3) and a positive correlation between gene expression and CG methylated genes (Fig. S6 and SI Text, section IV.2), as seen before in other insect species (28–33). Notably, however, DEGs tended to be hypomethylated in both species (Fig. 3E), and unlike the case in brain methylomes of adult honey bees (24, 34, 35), we found no evidence that phenotypes were associated with differentially methylated genes in our two species (t test, $P > 0.05$; SI Text, section IV.8). Analyses of alternative splicing revealed only 28 phenotype-specific isoforms expressed in *D. quadricaps* and none in *P. canadensis* (SI Text, section IV.9 and Dataset S3). This similarity between phenotypes is likely due to the global tendency of these species to express all isoforms simultaneously (Fig. 3F). Similar to DEGs, alternatively spliced genes (ASGs) were also hypomethylated compared with non-ASGs (Fig. 3E). This result may limit the role of DNA methylation in regulating phenotype-associated gene expression or

alternative splicing in our species, and contrasts with what has been described in the honey bee (26, 35–39).

MicroRNAs Are Not Preferentially Targeting Differentially Expressed Genes. Species-specific miRNA libraries were constructed from pools of individuals to include each phenotype to determine whether large numbers of miRNAs are shared between hymenopterans to the exclusion of the other insects and to identify potential *cis*-regulatory elements of DEGs. From our miRNA libraries, we identified 159 miRNA families (73 in *P. canadensis* and 86 in *D. quadricaps*), including 15 previously undescribed families (Fig. S7; SI Text, section V; and Dataset S4). We identified four families that are unique to hymenopterans and an additional nine families that were shared by apocritans to the exclusion of *Nasonia* and other insects. We found that miRNAs (40) were not preferentially targeting phenotype-specific DEGs, because although some DEGs appeared to be highly targeted, others were not (Dataset S4). Further work is needed to investigate miRNA expression levels in large numbers of individual queens and workers to rule out a role for miRNAs in caste differentiation.

Role for Conserved Toolkit Genes and Taxon-Restricted Genes in Regulatory Networks. Despite the low numbers of DEGs, we found evidence that DEGs were nonrandomly organized at the network level in both species. Weighted gene correlation network analyses identify groups of genes that covary significantly in expression as “modules” (41). These analyses identified 31 and 41 gene coexpression networks for the ant and wasp, respectively (SI Text, section III.5 and Dataset S5). DEGs were clustered nonrandomly across networks in both species (Fig. 4A). Only three (10%) and two (5%) network modules showed significant overrepresentation of DEGs in the ant and wasp, respectively, and only one network module in the ant showed evidence of functional enrichment for ribosomal terms (SI Text, section III.5). Phenotype-specific transcription in both species is therefore governed by subtle but coordinated coexpression networks.

There is a debate over the relative roles for core sets of conserved genes (42–48) and taxon-restricted genes (TRGs) (5, 44, 47, 49, 50) in the evolution of convergent phenotypes (7, 44, 46). We found evidence that both types of gene classes play peripheral roles in the molecular networks associated with phenotypic differentiation in our study species. In each species, we identified both classes of genes among DEGs, determined whether their functions were conserved, and ascertained their putative importance in the gene networks associated with phenotypic differentiation. There were significant levels of overlap in the identity of DEGs between the two species (reciprocal BLAST hits of DEGs; $n = 11$ genes, $P < 0.003$ relative to chance for both species; SI Text, section III.3 and Dataset S2), suggesting they are homologs. Some of these genes were the same as those genes that had been previously identified as conserved “toolkit” genes for alternative phenotypes in eusocial insects [e.g., *cytochrome*

Table 1. Transcriptome analyses

Species (no. of genes analyzed)	DEGs	Caste	NOISeq	EdgeR	Combined DEGs (% of total DEGs)	TRGs (% of DEGs)	Putative noncoding
<i>P. canadensis</i> (16,997)	67	Reproductive	56	36	64 (95)	5 (7.8)	2
		Nonreproductive	0	3	3 (5)	0	0
<i>D. quadricaps</i> (16,503)	147	Reproductive	29	55	74 (50.4)	9 (12.2)	2
		Nonreproductive	2	74	73 (49.6)	7 (9.6)	3

Two statistical methods were used to detect DEGs. EdgeR (a parametric approach) recognizes significant differences in gene expression when there is a large difference between the means of the groups. NOISeq-BIO (a nonparametric approach) tolerates much lower fold differences in gene expression between groups (as long as the ranges of the two conditions have little overlap) or if one of the groups shows evidence of high gene expression variation (“noisy” gene expression). The numbers of genes identified using EdgeR and NOISeq correspond to genes that were significantly up-regulated in reproductives or nonreproductives, as indicated. The number (and percentage as function of total DEGs) of TRGs and their coding potential are also given (full data are available in Dataset S2). Combined DEGs correspond to the union of genes detected by both EdgeR and NOISeq methods (note that some genes were detected by both methods).

Table 2. Methylation analyses

Species	Context	Total 1-kb regions analyzed	Global methylation, %	No. of HMRs (% of total 1 kb)	Methylated genes	HMR gene enrichment
<i>P. canadensis</i>	CG	172,660	2.79	1,060 (0.6)	731	1.8-fold
	Non-CG	192,001	3.62	1,057 (0.6)	314	1-fold
<i>D. quadriceps</i>	CG	244,626	3.01	7,065 (2.9)	4,360	1.8-fold
	Non-CG	245,493	1.26	12 (0.0)	6	0.7-fold
<i>A. mellifera</i> (24)	CG	194,707	2.29	8,046 (4.1)	3,861	2.7-fold
	Non-CG	212,728	0.11	0 (0.0)	0	0-fold

A minimum of 10% methylation level per 1-kb probe was used as a threshold to identify highly methylated regions (HMRs) in our two study species and in the honey bee *A. mellifera* as a comparison. Genes were defined as methylated if at least one HMR overlapped with a gene body (S/ Text, section IV.2; full data are available in Dataset S3).

P450, *vitellogenin*, *hexamerin-2*, *kruppel homolog 1* (42–48)], but others were not (e.g., *fibrillin-like* gene; *glutaminase*, *esterase*, and *myrosinase* enzymes; a gene coding for a lysozyme). Gene identity may be conserved, but not the direction of expression (5, 7): Four of 11 genes were worker-biased in the ant, whereas all 11 were queen-biased in the wasp (Dataset S2). Finally, conserved DEGs were not generally highly connected in the coexpression networks of either species (Fig. 4 C–F). This observation contrasts with eusocial insect species with phenotypes that are determined irreversibly during development, where conserved genes can play central roles in gene networks (44).

TRGs (those genes having no significant homologs in available genomic databases) were detected in DEG sets in both species (ant: 10%, $n = 16$; wasp: 7.5%, $n = 5$) (Fig. 4 C–F, Table 1, and Fig. S8) and at similar levels to TRGs across the whole genome

[ant (11.6% TRGs): $\chi^2 = 0.11$, $P = 0.74$; wasp (9.1% TRGs): $\chi^2 = 0.52$, $P = 0.47$; Dataset S5]. Taxon-restricted DEGs are likely to be new genes (short in length relative to annotated/known genes) (49) (Fig. S8) with unknown/novel functions (“guilt-by-association” network analysis) (41), because their nearest neighbors were also taxon-restricted (unknown function) (mean = 2.3 of the 10 most connected genes had BLAST hits; Fig. S8 and Dataset S5). Finally, taxon-restricted DEGs had similar low levels of connectivity to conserved genes in the networks of both species (generalized linear model) (ant: binomial errors, $P = 0.89$; wasp: quasibinomial errors, $P = 0.96$; Fig. 4 C–F), suggesting that conserved genes and previously unidentified TRGs are similarly important in phenotypic differentiation in these two species.

These data support the emerging hypothesis that conserved genes, new genes, and/or new regulatory networks are important

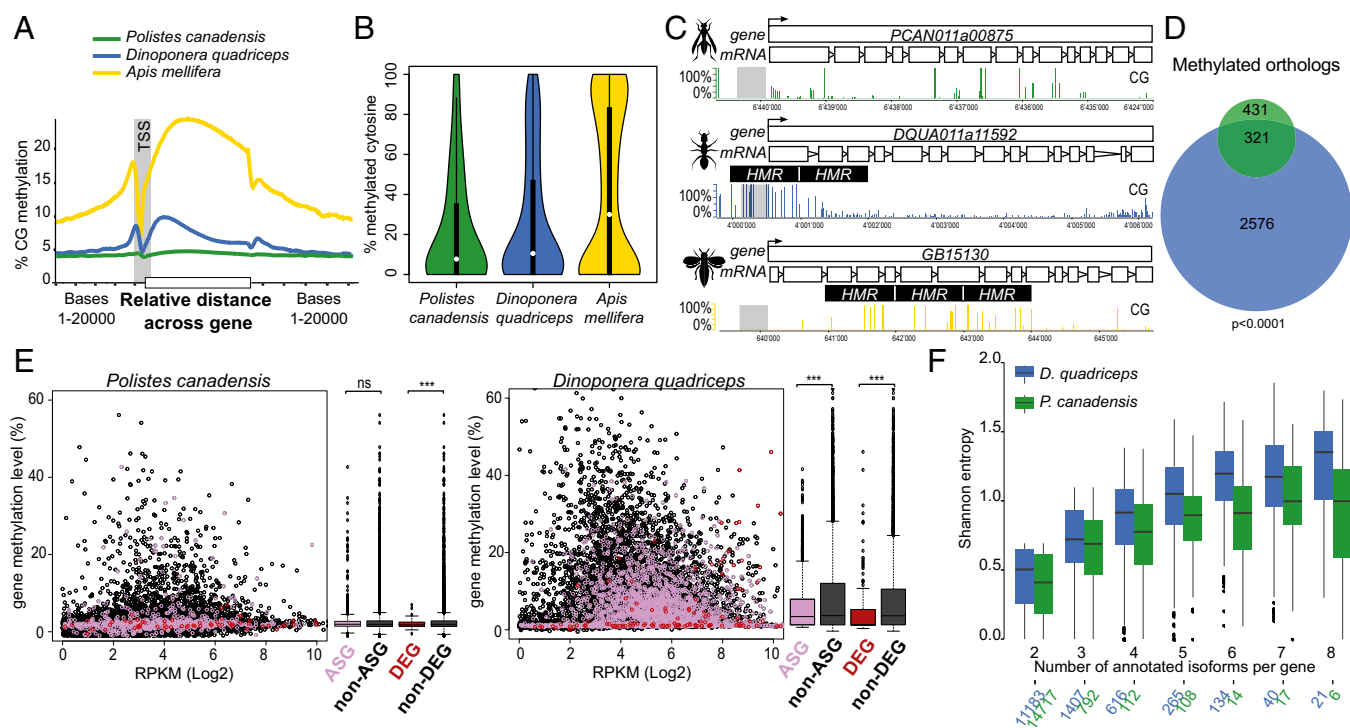


Fig. 3. Absence of distinct DNA methylation patterning. (A) Average CG methylation level in brain tissue along gene bodies and 20 kb of adjacent sequence for *P. canadensis* (green), *D. quadriceps* (blue), and *A. mellifera* (yellow). (B) Proportion of methylated cytosines within highly methylated regions (HMRs). Hartigan's dip test for unimodality: $D = 0.0184$ in *P. canadensis*, $D = 0.0257$ in *D. quadriceps*, and $D = 0.0849$ in *A. mellifera* ($P < 0.0001$ in all three species). (C) Screen shot from SeqMonk software showing the distribution of CG methylation in an orthologous gene in each of the three species. (D) Venn diagram of methylated orthologs: 74.5% (321 of 431) of the methylated genes in *P. canadensis* (green) overlap with *D. quadriceps* (blue). (E) Methylation distribution and summary box plots of the DEGs, ASGs, and non-ASGs, tested with Welch two-sample t tests. ns, not significant; RPKM, reads per kilobase per million. *** $P < 0.0001$. (F) Splicing entropy of annotated transcript isoforms. Shannon entropy grows with the number of annotated isoforms and with their equifrequency (entropy is 0 when only one isoform is expressed and high when all isoforms are expressed equally, Welch two-sample t test).

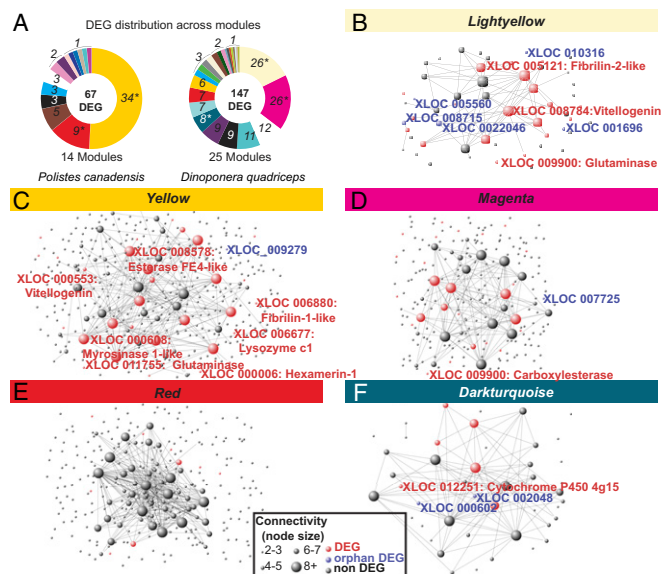


Fig. 4. Coordinated transcriptional network organization. (A) DEGs are nonrandomly distributed across modules (groups of genes with similar levels of expression): 14 of 41 DEGs in *P. canadensis* modules [binomial generalized linear model (glm) \times 2[13] = 162; $P < 0.0001$] and 25 of 31 DEGs in *D. quadricaps* modules (binomial glm \times 2[24] = 288; $P < 0.00001$). Colors correspond to the different modules. An asterisk indicates the modules that correlate significantly with phenotype. (B–F) Network graphs show the connectivity of annotated genes and TRGs in the modules that correlate significantly with phenotype. There were two modules in *P. canadensis* [yellow module, $P = 2.4 \times 10^{-23}$ (C); red module, $P = 14.1 \times 10^{-22}$ (E)] and three in *D. quadricaps* [light yellow module, $P = 9 \times 10^{-19}$ (B); magenta module, $P = 2.7 \times 10^{-42}$ (D); dark turquoise module, $P = 8.6 \times 10^{-4}$ (F)]. DEG fold enrichment in module: yellow (9-fold), red (3.6-fold), light yellow (21.5-fold), magenta (5.4-fold), and dark turquoise (7.7-fold). Nodes represent individual genes (with their XLOC gene name given). Edges indicate high coexpression between genes; edges with a correlation below specific thresholds are removed to aid visualization (41) [Thresholds: 0.27–1 (B), 0.31–1 (C), 0.15–1 (D), 0.24–1 (E), 0.12–1 (F)]. Connectivity (number of edges per node above the threshold) is indicated by node size. Annotated DEGs that are hubs [hubs defined as highly connected genes with more than five connections ($c > 5$)] are shown in red, and taxon-restricted DEGs that are hubs ($c > 5$) are shown in blue. Toolkit genes and TRG names are highlighted. Three genes that are DEGs in both species were found to be hubs in some networks: *myosinase* ($c = 16$) in *P. canadensis* and *vitellogenin* ($c = 14$) and *fibrillin* ($c = 8$) in *D. quadricaps*.

in the evolution of phenotypic diversity (5, 44, 47–51). Our analyses add to this hypothesis by identifying roles for both conserved genes and TRGs in highly plastic phenotypes.

Summary and Conclusions

We sequenced the genomes, miRNAs, multiple brain transcriptomes, and methylomes from two eusocial insect species whose life cycles depend on high phenotypic plasticity throughout life. This data includes the first aculeate wasp genome sequence to our knowledge. Both species displayed three key molecular signatures that may be molecular hallmarks for highly plastic phenotypes in simple eusocial insects. These key molecular signatures are as follows: (i) little molecular differentiation between phenotypes in transcription but subtle nonrandom differentiation at the transcriptional network level; (ii) no evidence of a role for DNA methylation or miRNAs in regulating phenotypic differentiation and an overall lack of distinct methylome patterning, together with evidence of methylation turnover; and

(iii) a similar role for both conserved toolkit genes and previously unidentified taxonomically restricted genes in phenotypic differentiation. These characteristics may allow plasticity in the regulation of the genome, and thus facilitate plasticity at the phenotypic level (52). The sequencing of more species with different levels of plasticity and multiple phenotypes will be required to confirm this hypothesis (6). However, the available data suggest that these hallmarks contrast with those hallmarks of eusocial insects with low plasticity like the honey bee and most ants, where a large proportion of genes, functionality, and network differentiation are associated with phenotypic differentiation (44, 53–58), and where phenotypes appear to be regulated by DNA methylation (24, 25, 30, 34, 35, 37, 59–62). Comparisons of species with contrasting evolutionary histories, as in our study species, will be especially valuable in revealing the molecular signatures at the origin of social evolution (e.g., in *P. canadensis*) and in reversions from complex to simple behaviors (e.g., in *D. quadricaps*). Methylome data from the brains of other ant (or wasp) species are not currently available. However, whole-body analyses of two species of ants revealed less defined methylome patterning and fewer differentially methylated genes between reproductive and nonreproductive phenotypes in *Harpegnathos* (high phenotypic plasticity) compared with *Camponotus* (lower phenotypic plasticity) (30), in support of our hypothesis. These insights, and the generation of the deep, multifaceted genomic resources for two model organisms with simple societies, help to plug a fundamental gap in our understanding of the molecular basis of phenotypic plasticity and serve to generate novel and important hypotheses on eusocial evolution. A particular focus for future work would be on whether the intriguing lack of coherent DNA methylation patterning and a key member of the enzymatic machinery (DNMT3) as regulators of alternative phenotypes is of general importance in permitting genomes to be highly responsive, as we have seen at the phenotypic level in social species with high phenotypic plasticity.

Methods

Detailed methodology and supporting information on sample collection (SI Text, section I), genome sequencing (SI Text, section II), RNA-sequencing (SI Text, section III), BS-seq (SI Text, section IV), and miRNA sequencing (SI Text, section V) are provided. A dataset (Datasets S1–S5) for every section is also provided.

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